

AperTO - Archivio Istituzionale Open Access dell'Università di Torino

**Multiple guests in a single host: interactions across symbiotic and phytopathogenic bacteria in phloem-feeding vectors - a review**

**This is the author's manuscript**

*Original Citation:*

*Availability:*

This version is available <http://hdl.handle.net/2318/1695049> since 2021-03-11T14:10:09Z

*Published version:*

DOI:10.1111/eea.12766

*Terms of use:*

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)

1

2

31 Multiple guests in a single host: interactions across symbiotic and phytopathogenic bacteria in

4

52 phloem-feeding vectors

6

7

83

9

10

114 Elena Gonella<sup>1\*</sup>, Rosemarie Tedeschi<sup>1</sup>, Elena Crotti<sup>2</sup>, Alberto Alma<sup>1</sup>

12

13

145

15

16

176 <sup>1</sup>Dipartimento di Scienze Agrarie, Forestali e Alimentari (DISAFA), Università degli Studi di Torino,

18

19

207 Grugliasco, Italy.

21

22

238 <sup>2</sup>Dipartimento di Scienze per gli Alimenti, la Nutrizione e l’Ambiente (DeFENS), Università degli

24

25

269 Studi di Milano, Milano, Italy.

27

2810

29

30

3111 \*Corresponding author: Elena Gonella, Dipartimento di Scienze Agrarie, Forestali e Alimentari

32

3312 (DISAFA), Università degli Studi di Torino, largo P. Braccini 2 10095 Grugliasco (TO) Italy. E-mail

34

3513 elena.gonella@unito.it

36

37

3814

39

40

4115 **Running title:** Bacterial interactions in phloem phytopathogen vectors

42

43

4416

45

46

4717

48

49

50

51

52

53

54

55

56

57

58

59

60

## Abstract

**Phloem-limited bacteria** are a major threat for worldwide agriculture due to the heavy economic losses caused to many high-value crops. These disease agents, namely phytoplasmas, spiroplasmas, liberibacters and *Arsenophonus*-like bacteria, are transmitted from plant to plant by phloem-feeding Hemiptera vectors. The associations established among pathogens and vectors often derive from co-evolution, and hence could result in a complex network of interactions involving also the whole microbial community harboured by the insect host. Interactions among bacteria may be beneficial, competitive, or detrimental for the involved microorganisms, and can dramatically affect the insect vector competence and consequently the spread of diseases. Interferences are observed among different pathogen strains competing to invade the same vector specimen, causing selective acquisition or transmission. Bacterial symbionts are another pivotal element for interactions **existing between vectors and phytopathogens**, because of their central roles for insect life cycle. Some symbionts, either obligate or facultative, were shown to have antagonistic effects on the colonization by plant pathogens, by producing or stimulating **the** insect production of antimicrobial substances, or competing for host infection. In other cases, evidences of mutual exclusion between symbiont and pathogen suggested possible detrimental influence on phytopathogens displayed by symbiotic bacteria; conversely examples of microbes enhancing pathogen load are available as well. Whether and how bacterial exchanges occurring in vectors affect the relationships between insects, plants and phytopathogens are still incompletely characterized issues, leaving room for many open questions concerning the significance of some traits of these multitrophic interactions. However, such complex interplays may have a serious impact on pathogen spread and control, having the potential to drive new strategies for the containment of important diseases.

**Keywords:** phytoplasma, liberibacter, spiroplasma, *Arsenophonus*, symbiotic bacteria, antagonism, competition

## 43 Introduction

44 Phloem-limited bacterial phytopathogens, which are among the most devastating agricultural threats  
 45 globally due to their wide host range and symptom severity, strictly rely on insect vectors to be spread  
 46 from plant to plant. These pathogenic bacteria are walled Proteobacteria ( $\alpha$ - and  $\gamma$ - subclades), and  
 47 wall-less Mollicutes. The first group encompasses the  $\alpha$ -proteobacterial ‘*Candidatus Liberibacter*  
 48 spp.’, including important pathogens of citrus and vegetable crops (Haapalainen, 2014), and two  
 49 *Arsenophonus*-related  $\gamma$ -Proteobacteria, namely ‘*Ca. Phlomobacter fragariae*’ and ‘*Ca.*  
 50 *Arsenophonus phytopathogenicus*’ (Bressan, 2014). Plant pathogenic Mollicutes embrace the genera  
 51 ‘*Ca. Phytoplasma*’ and *Spiroplasma*.

52 All vectors of plant pathogenic bacteria residing in the phloem are Hemiptera belonging to the  
 53 suborders Auchenorrhyncha (with the families Cixiidae, Dictyopharidae and Flatidae in the  
 54 Fulgoromorpha infraorder and Cicadellidae in the Cicadomorpha infraorder) and Sternorrhyncha  
 55 (superfamily Psylloidea). Vectors are able to ingest bacteria by feeding in the phloem with their  
 56 piecing-sucking mouthparts. Liberibacters are transmitted by psyllids, and *Arsenophonus*-like  
 57 bacteria are vectored by planthoppers in the family Cixiidae. On the other hand, phytoplasmas are  
 58 transmitted by leafhoppers (family Cicadellidae), planthoppers (superfamily Fulgoroidea), and  
 59 psyllids (superfamily Psylloidea); while spiroplasmas are vectored by leafhoppers only (Gasparich,  
 60 2010).

61 The interactions between plant pathogens and their vectors are not limited to a carrier-carried relation:  
 62 different species or strains of a plant pathogen have divergent behaviours in different insect hosts.  
 63 Moreover, phytopathogenic bacteria are included in a complex network of interactions occurring in  
 64 vectors, being actual members of the multifaceted insect microbiomes, which have a significant  
 65 influence on the biology of the hosts. Members of the Hemiptera, including all of the vectors of  
 66 phloem-limited bacterial plant pathogens, rely on bacterial symbionts for supplying nutrients lacking  
 67 in their unbalanced diet (Baumann, 2005). The nutritional provisioning operated by obligate

symbionts has been a crucial condition for insect persistence and diversification on a limited food niche such as plant phloem (Skidmore and Hansen, 2017), then affecting the host range of vectors. High polyphagy deriving from mutualistic associations may in turn influence the chance of different plants to be infected by a plant pathogen. Moreover, facultative symbionts are commonly found in many vectors, showing protective functions, or being capable to manipulate the host's reproduction (Zchori-Fein & Bourtzis, 2011). In addition, different species or strains of plant pathogens may be hosted by the same individual vector (Table 1), possibly being transferred together to the host plant (Bosco & D'Amelio, 2010). Such multipartite interactions most commonly result in microbial synergies or interference, with potential implications for bacterial transmission as well (Bosco & D'Amelio, 2010; Saldaña et al., 2017). This review summarizes the available knowledge concerning microbial exchanges occurring in the vectors of phloem bacterial pathogens, with special regard to the consequences on their transmission. Disease management could take advantage of these interactions to develop microbe-based control strategies (Crotti et al., 2012) (Figure 1). Indeed, despite their capability to easily adapt to, and grow in, different hosts such as plants and insects, currently these phloem-restricted bacteria cannot be cultured or are difficultly cultivated in cell-free media –with few exceptions such as spiroplasmas and a single liberibacter species– (Perilla-Herao & Casteel, 2016), thus limiting experimentations aimed to identify new control strategies. Control is generally based on the use of healthy plant propagation material, elimination of symptomatic plants, and control of insect populations spreading the disease. Unravelling the interactions established between phytopathogens and insect symbionts could offer an interesting tool to impair the transmission of phloem-limited plant pathogens in a sustainable perspective.

## Phloem-limited bacterial plant pathogens

### Liberibacters

Transmitted by psyllids, 'Ca. Liberibacter' pathogens include primarily obligate parasites of plants and insects, responsible for several plant diseases, among which huanglongbing (HLB) in citrus trees

and zebra chip (ZC) in potatoes are the most severe ones in terms of crop damage and economic losses (Gottwald et al., 2007; Haapalainen, 2014). Three species of ‘*Ca. Liberibacter*’ have been indicated as the causal agents of citrus HLB, previously known as citrus greening, i.e. ‘*Ca. L. asiaticus*’ (CLas), ‘*Ca. L. africanus*’ (CLaf), and ‘*Ca. L. americanus*’ (CLam), the names of which have been derived from the continents where these bacteria have been originally found and are mainly distributed (Haapalainen, 2014). While CLaf is transmitted by the African citrus psyllid *Trioza erytreae* Del Guercio (McClellan & Oberholzer, 1965), CLas and CLam are mainly vectored by the Asian citrus psyllid *Diaphorina citri* Kuwayama (Capoor et al, 1967; Teixeira et al., 2005). *D. citri*, native to southeastern Asia, has been recently diffused in America probably in consequence of international commerce (Halbert & Núñez, 2004; Bayles et al., 2017). Despite similar symptoms are recorded after infection by each of the three HLB-causing species, CLas is the most destructive one, inducing devastating epidemics in several countries (Haapalainen, 2014). On the other hand, ZC in potatoes and other diseases in vegetable crops are caused by ‘*Ca. L. solanacearum*’ (CLso), which has been initially indicated with the name ‘*Ca. L. psyllauros*’ (Liefting et al., 2009). Geographically distinct CLso haplotypes are known, whose differential distribution results in the association with separate plant and insect host species. While in North America and Oceania this pathogen is vectored by the potato/tomato psyllid *Bactericera cockerelli* Šulc, causing severe damage in potato and tomato crops, in Europe -where it is transmitted by psyllids of the species *Trioza apicalis* Förster and *Bactericera trigonica* Hodgkinson- it is associated with diseases of the Apiaceae family plants, such as carrot and celery.

In the last years, other liberibacter species have been identified, i.e. ‘*Ca. L. europeus*’ (CLEu) and *Liberibacter crescens*, but differently from the aforementioned species these latter are not reported as phytopathogens, rather showing an endophytic behaviour (Raddadi et al., 2011; Leonard et al., 2012). Interestingly, *L. crescens*, found in mountain papaya in Puerto Rico, can be grown in axenic cultures, making it an optimal candidate to study liberibacters’ biology (Leonard et al., 2012; Fagen

et al., 2014a,b). On the other hand, CLeu, reported as an endophyte of pear, apple, blackthorn and hawthorn, transmitted by *Cacopsylla* spp. (Raddadi et al., 2011; Camerota et al., 2012), has been recently indicated as a pathogen in Scotch broom (*Cytisus scoparius*) in New Zealand (Thompson et al., 2013). Recently, other two new candidate liberibacter species were recently reported: the ‘*Ca. Liberibacter caribbeanus*’ (CLca) detected in *Citrus sinensis* (L.) Osbeck and in the citrus psyllid *D. citri* from Colombia (Keremane et al., 2015) and the ‘*Ca. Liberibacter brunswickensis*’ (CLbr) detected in the native Australian eggplant psyllid, *Acizzia solanicola* Kent & Taylor (Morris et al., 2017). Neither these new species were associated with plant disease but a co-evolution with psyllids as secondary symbionts is inferred (Morris et al., 2017).

#### *Arsenophonus*-like bacteria

*Arsenophonus* genus includes not only plant pathogens, but also insect parasites and symbionts (Bressan, 2014). For instance, in a survey performed on 136 arthropod species it has been found that *Arsenophonus* bacteria are associated with 5% of the tested hosts (Duron et al., 2008), where they can establish complex interactions with beneficial or parasitic features (Wilkes et al., 2011). Conversely, two species are cause of disease to strawberry and sugar beet plants (Danet et al., 2003; Bressan et al., 2008). The first pathogenic agent was discovered at the end of last century in France on strawberries affected by marginal chlorosis. Because at that time very little was known about this genus, the pathogen was considered as a separate species that was named ‘*Ca. Phlomobacter fragariae*’ (Zreik et al., 1998); nonetheless the increase of sequence data availability led to propose it to be an *Arsenophonus* (Bressan, 2014). The other plant pathogenic *Arsenophonus* is ‘*Ca. Arsenophonus phytopathogenicus*’ which infects sugar beet, causing a disease defined as “basses richesses” syndrome, because diseased plants show decreased sugar content (Richard-Molard et al., 1995). The insect vectors of pathogens in the *Arsenophonus* group are cixiids: ‘*Ca. Phlomobacter fragariae*’ is vectored by *Cixius wagneri* (China) (Danet et al., 2003), whereas ‘*Ca. Arsenophonus phytopathogenicus*’ is transmitted by *Pentastiridius leporinus* (L.) (Gatineau et al., 2002). These two



pathogens are phylogenetically distinct, and can differentially interact with plants and insects in different contexts. '*Ca. Arsenophonus phytopathogenicus*' was observed in Italy to be related to a strawberry marginal chlorosis disease (Terlizzi et al., 2007); likewise it was detected in *C. wagneri*, which was able to inoculate it to sugar beet plants, whereas strawberries were not infected (Bressan et al., 2008). Moreover, the epidemiology of this group of diseases is complicated by the fact that they can be induced also by phytoplasmas transmitted by *Hyalesthes obsoletus* Signoret (Gatineau et al., 2002, Danet et al., 2003). Even being plant pathogens, there is evidence that many traits of *Arsenophonus*-like bacteria are characteristic of an insect symbiont lifestyle, such as reproductive tissue colonization and vertical transmission, absence of entomopathogenic activity, high infection rate and a life cycle prevalently related to insect hosts (Bressan, 2009b; 2014). Thus, these bacteria could easily initiate new associations with additional cixiid species. The complexity of their associations with insects and plants, jointly to cixiids' capability to easily adapt to new environments and host plants, could effectively explain the increasing appearance of emerging *Arsenophonus*-related diseases.

### Phytoplasmas

Phytoplasmas are known to be responsible for diseases in over a thousand of economically important crops globally distributed (Marcone, 2014): typical symptoms include yellowing, witches' broom, virescence, phyllody, bolting, reddening of leaves and stems, decline and stunting of plants (Hogenhout et al., 2008). To date, all known phytoplasmas are reported to be pathogenic for at least one plant, even though asymptomatic hosts may be recruited. Phytoplasma taxonomy has been hampered by their recalcitrance to be cultured in vitro; therefore these bacteria are partially classified in the provisional genus '*Ca. Phytoplasma*' based on sequence analysis; up to now 42 '*Ca. Phytoplasma*' species have been reported (Zhao & Davis, 2016). A more exhaustive categorization defines phylogenetic clusters (16SrI-XXXIII groups, each one divided in many subgroups) according to 16S rRNA gene sequence (Lee et al., 1993, 1998b; Zhao & Davis, 2016).



Since most phytoplasmas are capable to cause symptoms to a number of plants belonging to different families, such phytopathogens are regarded as some of the most troubling disease agents in these areas. Moreover, some phytoplasmas are successfully transmitted by polyphagous vectors, furtherly incrementing their chance to infect a huge number of plants. For example, Aster Yellows phytoplasmas (16SrI) are vectored by many polyphagous leafhoppers to several plants (Weintraub & Beanland, 2006), including different flowers, vegetables, or grapevine. The broad range of wild and cultivated plants that are affected by these pathogens can be explained by the polyphagy recorded for most of vectors, along with the great diversity of phytoplasma subclades within this group (Hogenhout et al., 2008).

Considering vector-phytoplasma interplays, many specific interactions are acknowledged between different phytoplasma phylogenetic groups and distinct taxa of vectors. As an example, only leafhoppers in the family Cicadellidae have been reported to transmit phytoplasmas of the 16SrI group (Alma et al., 2015). On the other hand, many phytoplasmas are indistinctively vectored by distant insects. For instance, phytoplasmas of the phylogenetic groups 16SrV and 16SrXII may be vectored by members of either Fulgoromorpha and Cicadomorpha, and 16SX phytoplasma can be transmitted both by Auchenorrhyncha and Sternorrhyncha (Alma et al., 2015). However, a single family with major vector importance can be generally recognized even for pathogens transmitted by distinct taxa: in the case of 16SrV phytoplasmas, most of vectors belong to Cicadellidae, 16SrXII phytoplasmas are mainly transmitted by cixiids, and the major vectors 16SrX phytoplasmas are members of Psyllidae.

### Spiroplasmas

Spiroplasmas are regarded as an extremely harmful group for global agriculture, even though only few species have been accounted as phytopathogens, i.e. *Spiroplasma citri* in citrus, *S. kunkelii* in maize and *S. phoeniceum* in aster (Gasparich, 2010). All plant pathogenic spiroplasmas are phylogenetically related, being included in the same taxonomic lineage, namely the Citri clade

(Garsparich, 2010). Despite spiroplasmas and phytoplasmas establish similar pathogenic relationships with host plants, inducing analogous symptoms, major biological differences are evident between these genera. Distinctions include the bacterial shape, as spiroplasmas are characterized by the helical morphology and phytoplasmas are pleomorphic, and cultivation suitability, as spiroplasmas can be cultured in nutrient-rich media while phytoplasmas are recalcitrant to cultivation (Gasparich, 2010).

*S. citri* is mainly related to heavy economic losses to citrus productions; however this pathogen, as well as its vectors, may be found on many different host plants. Namely, *S. citri* is the agent of citrus stubborn, brittle root disease of horseradish, sesame yellowing, and carrot purple leaf (Zarei et al., 2017); it is transmitted by the leafhoppers *Circulifer haematoceps* (Mulsant & Rey) in the Mediterranean basin and *Circulifer tenellus* (Baker) in North America (Renaudin, 2006). The main areas affected by *S. citri*-related diseases are the Mediterranean countries of Europe, North Africa, and western Asia, as well as the Nearctic region, whereas the pathogen is absent in South America.

*S. kunkelii* is an important pathogen of maize crops, even though its distribution is restricted to the Americas. Its natural vector is the cicadellid *Dalbulus maidis* (DeLong & Wolcott), which is a specialist of the genus *Zea* present in the Nearctic and Neotropical areas. *D. maidis* is co-evolved with maize, where it can be among the most prevalent leafhoppers (Palomera et al., 2012).

The third plant pathogenic spiroplasma species is *S. phoenicium*, which was retrieved from periwinkle plants affected by yellows in Syria. This pathogen is experimentally transmitted by the leafhopper *Macrostes fasciifrons* (Stål); however, at present no information is available concerning the natural vectors of *S. phoeniceum* in the infested area (Saillard et al., 1987).

### **Bacterial phytopathogen-vector relations**

In the vectors, the phloem-restricted pathogens are transmitted in a persistent manner: once ingested by trophic activity on infected plants, bacterial cells multiply in the insect midgut, cross the

epithelium, replicate in the hemolymph and, ultimately, infect the salivary glands to be further injected in the new host plant (Figure 1; Gasparich, 2010; Bressan, 2014; Haapalainen, 2014). This process implies complex interplays, spanning from beneficial to adverse. A benign role was suggested for CLas in *D. citri* (Duan et al., 2009; Mann et al., 2011), although an increased susceptibility to selected insecticides was observed in infected psyllids, resulting in fitness decrement (Mann et al., 2011). Similarly, a negative density-dependent effect of CLso infection on the fecundity of *B. cockerelli* was reported by (Nachappa et al., 2014), whereas no significant detrimental effects on the biology of infected individuals occurs according to Thinakaran et al (2015). Effects of vector manipulation by a phytopathogen have been observed also at the hemolymph level, as in CLas-infected *D. citri* showing changes in proteins related to energy metabolism, immunity, and lipid transport (Kruse et al., 2018). Differential effects have been reported for insect-phytoplasma associations: for example, shorter survival and a lower egg production were observed in individuals of *Scaphoideus titanus* Ball infected by 16SrV phytoplasmas (Bressan et al., 2005a), whereas a positive influence have been recorded for 16SrI phytoplasmas in *Macrosteles quadrilineatus* DeLong & Caldwell (Beanland et al., 2000).

The molecular mechanisms regulating plant pathogens retention, multiplication and spread in the body of some species, and not in others, are still poorly understood. The biological adaptation of vectors to harbour plant pathogens suggest a co-evolution between insects and bacteria; however, these interactions have polyphyletic traits, indicating multiple independent evolution events (Orlovskis et al., 2015). The evolution of pathogen transmission shares some traits with insect symbiosis, as most of plant pathogens are phylogenetically related to many symbiotic bacteria of Hemiptera, and similarly to endosymbionts they have reduced genomes, reflecting the adaptation to obligate associations (Bendix & Lewis, 2018). Indeed, a major consequence of a host-dependant life style is an extreme gene loss, due to the lack of a selection process capable to maintain superfluous genes in the rich environment provided by the insect body (Latorre & Manzano-Marín, 2017). In most

cases, the associations between plant pathogens and their vectors are believed to be originated from bacterial internalization and successful survival in insects feeding transiently in infected plants (plant-first model). Conversely, some phytopathogens, especially those in the Enterobacteriaceae family, may have been initially insect commensals (i.e. non-harmful associates) that have evolved as plant pathogens following repeated inoculations in the phloem by their insect hosts (insect-first model) (Bové & Garnier, 2002; Nadarasah & Stavrinides, 2011).

Traits affecting vector suitability and specificity are thought to be related to difference in insect physiology, immunity, and behaviour, as well as to their geographical and seasonal distribution (Perilla-Hernao & Casteel, 2016). For instance, divergent plant host-dependant feeding behaviours have been suggested to play an important role in differential transmission competence observed in the leafhopper phytoplasma vectors *Euscelidius variegatus* (Kirschbaum) and *Empoasca decipiens* Paoli (Galletto et al., 2011). Moreover, the vector immune system may limit pathogen invasion. In *D. citri*, CLas acquisition by adult specimens was proven to be significantly less efficient than by nymphs due to differential immune responses, like melanization and apoptosis of gut cells (Kruse et al., 2017). Similarly, immune response may be the cause of limited phytoplasma cell number found in non-transmitting individuals of different vector species after experimental exposure to the pathogens (Galletto et al., 2009). A crucial phase of the transmission process is the protein interaction between pathogen cells and those of the host, regulating pathogen crossing of gut and salivary glands epithelia. The main strategy for bacterial internalization reported for plant pathogenic agents is endo-exocytosis (Kwon et al., 1999; Hogenhout et al., 2008; Cicero et al., 2016), mediated by different membrane proteins (Labroussaa et al., 2010, 2011; Béven et al., 2012; Duret et al., 2014; Konnerth et al., 2016; Arricau-Bouvery et al., 2018). The absence of specific adhesion machinery to host cells seriously weakens the vector competence (Weintraub & Beanland, 2006). For example, *S. citri* strains lacking adhesion-related proteins are not transmissible by insects (Kruse et al., 2017).

The transmission of a plant pathogen by vectors is affected also by the fact that different species or strains of a plant pathogen have divergent behaviours in different insect hosts. This is especially observed for those phytopathogens that most probably derive from insect symbionts, such as *Arsenophonus* bacteria and spiroplasmas. Both the genera *Arsenophonus* and *Spiroplasma* encompass inter- and intracellular symbiotic bacteria displaying a diversity of roles, from mutualism to reproductive manipulation, or may even be entomopathogenic (Gasparich, 2010, Bressan, 2014). In ‘*Ca. A. phytopathogenicus*’ and ‘*Ca. P. fragariae*’, it has been shown that the exploitation of plants resulted from independent evolutionary events from a common endosymbiotic ancestor (Bressan, 2014). This evidence, along with the observation of typical symbiotic traits in insects, like high prevalence and maternal transmission, suggests their transition from endosymbiotic to plant pathogenic life style (Bressan, 2014). Besides, some species belonging to other phytopathogen groups could actually derive from insect commensals. For example, phylogenetic studies demonstrated a match between the affinity level of liberibacter species restricted to different continents and the geographical distribution of psyllid hosts. This supported the hypothesis of a co-evolution between CLbr, behaving as an insect secondary symbiont, and its host *A. solanicola* (Morris et al., 2017). On the other hand, co-evolved associations involving a plant pathogen and an insect vector may lead to mitigate possible harmful effects exhibited on the host fitness (Purcell, 1982). The growing number of observed transitions from insect endosymbiosis to plant pathogeny and vice versa is certainly indicative of the possibility that new bacterial species, currently believed to be horizontally transmitted insect commensals or mutualists, will become emerging plant pathogens in the future.

The study of phytopathogen-vector interactions has a remarkable pertinence from a disease containment perspective, because differential molecular targets for control could be derived from distinct associations involving co-evolution, mutualism or insect injury. For example, the enhancement of insect immunity could be a specific control objective in case of pathogen-vector interactions where the bacterium is definitely recognized and attacked by immune cells due to non-

beneficial interchange (Weiss & Aksoy, 2011). In contrast, some phytopathogens that are anciently related to and co-evolved with their insect hosts are able to escape the immune response. For example, *S. citri* has been reported to evade phagocytosis and limit phenoloxidase activity in its vector *C. haematocephus* (Eliautout et al., 2016). In those cases, control approaches based on immune augmentation may be insufficient.

### Multiple pathogen infections and competition

The interaction among pathogens, plants and vectors can be extremely complex. Mixed infections by different bacterial pathogens can quite commonly be observed in the phloem of the same plant. The simultaneous occurrence of multiple pathogens in the same plant is rather frequent in herbaceous plants and trees belonging to many families; either related and phylogenetically distant pathogenic agents may co-exist (Križanac et al., 2010; Nicolaisen et al., 2011; Arratia-Castro et al., 2016; Satta et al., 2016; Swisher et al., 2018). Moreover, a single insect can feed on several plants, or even different plant species, during its life cycle, possibly being exposed to mixed pathogen infections. As a consequence, insect vectors may acquire many pathogen species or strains during the same feeding event, or by feeding sequentially on host plants infected by different bacteria (Križanac et al., 2010; Raddadi et al., 2011; Swisher et al., 2018) (Table 1). However, in some cases, the co-occurrence of multiple pathogens in the same insect's body is inhibited by interferential interactions such as selective acquisition or transmission of a single microbe (Bosco & D'Amelio, 2010). For example, in the leafhopper *Dalbulus maidis* (Delong & Wolcott), which is the natural vector of maize bushy stunt phytoplasma (MBSP) and corn stunt spiroplasma (CSS), competition for transmission was reported after co-occurrence during a long-term latency period (de Oliveira et al., 2007). This competition resulted in suppression of prolonged transmission of MBSP after acquisition of CSS, as the latter is thought to have faster rates of multiplication and spread, hence being more competitive during the latency period required for successful transmission. Similar results were obtained with the cicadellid *M. quadrilineatus*, vector of several strains of Aster Yellows Phytoplasma. Leafhoppers



exposed to sequential acquisition of different phytoplasma strains most frequently transmitted the first provided isolate exclusively (Freitag, 1967). These evidences suggest competitive colonization of the insect's body, where the first strain starting multiplication and reaching the salivary glands is more competitive and hence preferentially transmitted (Bosco & D'Amelio, 2010). The same competitive colonization process was proposed for *Osbornellus horvathi* Matsumura, since 'Ca. P. asteris' and 'Ca. P. phoenicium' double-infected adult leafhoppers were able to transmit the former, but not the latter, to different plants in experimental conditions (Rizza et al., 2016). Considering *Arsenophonus*-related plant pathogens, no specific transmission trial from double-infected sources has been reported yet; however there are evidences that separated populations of *Cixius wagneri* (China), the only known vector of both pathogens, exclusively transmit 'Ca. A. phytopathogenicus' or 'Ca. P. fragariae' but do not carry the two bacteria together (Bressan et al., 2008). Many factors must be taken into account to explain exclusive pathogen acquisition by *C. wagneri*, including vector ecology and population dynamics, which could lead to limited chance for the same individual to be exposed to both pathogens; however the competition between 'Ca. A. phytopathogenicus' and 'Ca. P. fragariae' for insect colonization cannot be ruled out.

The competition between two bacterial pathogens in the vectors has been better dissected by Rashidi et al. (2014), by using the leafhopper *E. variegatus* and two unrelated phytoplasmas, namely Chrysanthemum Yellows phytoplasma (CYP) and Flavescence Dorée Phytoplasma (FDP), experimentally transmitted to broad bean plants. The authors found that insects sequentially exposed to acquisition of CYP and FDP showed unilateral interference, with the suppression of FDP transmission regardless of the feeding order. On the other hand, the acquisition of each pathogen was not affected by the presence of the other one, suggesting no competition at the earlier infection stages. The barrier where competition takes place was rather identified in salivary glands, which were more rapidly invaded by CYP due to its capability to multiply faster than FDP, even though the latter bloomed to higher concentrations. The higher speed in reaching salivary glands displayed by CYP



was suggested to be related to: i) long co-evolution with the insect host and consequent mitigated immune response, and ii) broad phytoplasma host range supporting the evolution of traits that promote acceptability by a broad vector range (Rashidi et al., 2014). Transcriptomic analysis of infected leafhoppers with single phytoplasma strains demonstrated the activation of insect immune response (by activation of Kazal type 1 serine protease inhibitor and melanisation pathway) after infection by FDP, which reduces the host fitness and is then perceived as a potential pathogen (Galletto et al., 2018). Instead, the most competitive CYP increased energy metabolism, providing molecular confirmation for different competition levels.

The knowledge on competition between co-occurring pathogen strains in the same host, although being still limited, could support the study of pathogen transmission. Indeed, the observation and characterization of competition events may contribute to unravel meaningful details of the processes determining insect invasion and spread of phytopathogens, possibly identifying weaknesses of single associations and revealing new control targets. Moreover, competitive transmission of different plant pathogens may seriously alter disease epidemiology in the field.

### Symbiont-pathogen interactions

The Auchenorrhyncha and Sternorrhyncha, including the vectors of plant pathogenic bacteria, harbour both obligate and facultative endosymbionts which play important roles in supplying nutrients and providing the host with other fitness benefits (Baumann, 2005; Morrow et al., 2017). The main obligate (primary) symbiont are ‘*Ca. Sulcia muelleri*’ in Auchenorrhyncha, and ‘*Ca. Carsonella ruddii*’ in psyllids. Moreover, *Sulcia* requires complementary (co-primary) symbiotic bacteria to integrate its nutrient supply to the insect (McCutcheon & Moran, 2010). Similarly, psyllids harbour secondary symbionts, such as *Sodalis* or *Arsenophonus* bacteria, with nutritional roles (Morrow et al., 2017). In addition, the function of some symbionts of hemipterans vectors is still unrecognized. For example, many bacteria generally known as reproductive manipulators, such as *Wolbachia*, *Cardinium*, *Rickettsia* and *Arsenophonus*, have been found in several vector species;

however their role has not been characterized yet (Marzorati et al., 2006; Gonella et al., 2011; Jing et al., 2014; Morrow et al., 2017; Iasur-Kruh et al., 2017). Moreover, some insect beneficial microorganisms (e.g. *Rickettsia* and *Cardinium*), capable to colonize the salivary glands, may be transferred from insect to plant and vice versa, possibly establishing endophytic relationships as well (Caspi-Fluger & Zchori-Fein, 2010; Gonella et al., 2015; Iasur-Kruh et al., 2017). Despite the emerging recognized need to study microbial communities affiliated to non-model insects (Prosdocimi et al., 2015), which recently led to a growing number of evidences of co-existence of plant pathogens and other microbes in the insect vectors, few studies directly investigated their interactions (Table 2). Symbiont-pathogen exchanges were firstly studied in psyllids, and more specifically in the CLas vector *D. citri*. This psyllid harbours three main endosymbionts: a species of *Wolbachia*, the  $\gamma$ -Proteobacterium '*Ca. Carsonella ruddii*', an endosymbiont which may provide nutritional benefits to its host (Thao et al., 2000), and '*Ca. Proffittella armatura*', a  $\beta$ -Proteobacterium with defensive function (Nakabachi et al. 2013). Fagen et al. (2012) firstly observed a negative correlation between CLas infection rate with the relative abundance, within the microbial community, of *Proffittella*. Based on its genome sequence, *Proffittella* was predicted to produce defensive toxins, i.e. diaphorin and diaphorin-related polyketides. CLas-infected [CLas(+)] insects were found to have dramatically elevated levels of two proteins involved in polyketide biosynthesis. In contrast, the protein responsible for initiating diaphorin biosynthesis is down-regulated in CLas(+) *D. citri* (Ramsey et al., 2015). Moreover, Ramsey et al. (2015) observed that the ratio between levels of diaphorin and the related polyketide is significantly increased in CLas (+) compared to CLas uninfected [CLas(-)] *D. citri*, suggesting changes in *Proffittella* polyketide metabolism in response to the presence of the pathogen or in direct or indirect response to changes induced by the pathogen in infected plants. The up-regulation of the polyketide synthase (PKS) gene expression in CLas(+) *D. citri* may be a specific response of *Proffittella* to the presence of CLas, as part of an infection response that may be mediated by *D. citri* (Ramsey et al., 2015). Such an interactive response may involve

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

391 *Carsonella* as well, which could provide the host with essential amino acids required for polyketide  
392 production (Ramsey et al., 2015).

393 Besides psyllid-liberibacter interactions, further evidences of antagonistic relationships between  
394 symbiotic bacteria and plant pathogens are reported for some Auchenorrhyncha vectors of  
395 phytoplasmas. A bacterium in the Xanthomonadaceae, provisionally named *Dyella*-like bacterium  
396 (DLB) (Iasur-Kruh et al., 2017), was isolated from the planthopper *H. obsoletus*, and showed anti-  
397 phytoplasmal activity in inoculated plants (Iasur-Kruh et al., 2018). Indeed, despite being isolated  
398 from an insect source, DLB showed endophytic traits: it was consistently found in the wild bush *Vitex*  
399 *agnus-castus* L., and it was able to long-term colonize the phloem of different plant species, including  
400 many hosts of phytoplasmas and liberibacters (Lidor et al., 2018). Once established in grapevines  
401 infected by phytoplasmas, DLB reduced disease symptoms (Iasur-Kruh et al., 2018). Based on DLB  
402 genome analysis, the authors suggested that such a drop of symptoms is related to inhibition of  
403 pathogens, rather than competition or production of substances stimulating plant growth or defense  
404 (Lahav et al., 2016; Iasur-Kruh et al., 2018). Moreover, DLB was demonstrated to inhibit the growth  
405 of the cultivable model Mollicute *Spiroplasma melliferum* (Iasur Kruh et al., 2017).

406 Acetic acid bacteria in the genus *Asaia* are widespread in insects, including leafhoppers transmitting  
407 phytoplasmas, and they were proposed to interact with insect vectors, possibly altering their spread  
408 (Crotti et al., 2009). Strains with different phenotypes previously isolated from mosquitoes were  
409 orally supplied to the experimental vector of FDP *E. variegatus*, which was successfully colonized.  
410 One *Asaia* strain producing an air-liquid interface biofilm, after establishing in *E. variegatus*, reduced  
411 its acquisition of FDP from broad beans in experimental conditions (Gonella et al., 2018). These  
412 authors suggested that the strain of *Asaia* could affect the capability of the phytoplasma to cross the  
413 gut epithelia for reaching salivary glands, even though the mechanisms regulating this interference  
414 remain to be elucidated. However, such an alteration was imperfect and, when the pathogen  
415 succeeded in colonizing the insect, transmission rates to broad beans were similar to those recorded  
416 for control leafhoppers unexposed to *Asaia* (Gonella et al., 2018).

Additional **interplays** between symbiotic bacteria and plant pathogens **have** been suggested by multiple prevalence studies, as in some **cases positive correlation or mutual exclusion could be detected** between symbiotic and phytopathogenic bacteria. For example, the obligate symbiont *Nasuia*, largely widespread in the family Cicadellidae, is present in most of leafhopper species transmitting phytoplasmas, while non-vector species were shown to lack it (Wangkeeree et al., 2012). It has been suggested that *Nasuia* could be required for successful transmission. Likewise, in the planthopper FDP vector, *Dictyophara europaea* L., a negative correlation between infections by phytoplasma and *Wolbachia* was reported, suggesting that the *Wolbachia* strain infecting *D. europaea* displays antagonistic activities against the pathogen, or alternatively competes for insect colonization (Krstić et al., 2018). On the other hand, in *D. citri* an increase in the ubiquitous *Wolbachia* titre was reported with CLas infection (Fagen et al., 2012), indicating a more complicated interplay mechanism with strain-specific variability. **Direct interaction has been documented between *Wolbachia* and CLas, as the first suppress the holing lytic promoter in a CLas-infecting phage in *D. citri* (Jain et al., 2017).**

**The studies regarding synergies and interferences between symbiotic agents and plant pathogens offer significant cues for disease treatment; moreover, further work is still required to describe new interactive associations. Future work concerning such interplays should be aimed not only to identify direct anti-pathogen activity expressed by symbionts, but also to alter the mutualistic exchange recorded among vectors, symbionts and phytopathogens, and to influence insect ecology (e.g. by driving plant choice and governing interactions with stresses).**

## Conclusions and open issues

The interactive roles of phytopathogenic and symbiotic bacteria in insects certainly represent an emerging topic for researchers focusing on the transmission process of disease agents. A multi-actor picture, involving insects, plants, and microbes, is resulting as the condition where the transmission

of plant pathogens arises. Consequently, the bacterial interactions occurring in insects affect the life cycle of the host as well. First, considering the reported evolutive bilateral transition of the role of many disease agents in their vectors from symbiotic to phytopathogenic life style, the effects of these bacteria are a key issue for the study of insect-microbe relationships; however they are still mostly unknown. Such effects may also result in the uneven competitive behaviours described for both closely and distantly related pathogens. Many questions arise from this hypothesis. How is insect immunity involved in differential growth rates of plant pathogens? What are the traits of vector-pathogen interaction originating possible diversity in host responses? Are these bacteria at different steps of transition from symbiont to pathogen or vice versa (e.g. the most competitive pathogens supply the host with fitness advantages)? Most of these questions were addressed by Galetto et al. (2018) using the *E. variegatus*-CYP-FDP model, but more work is needed to expand the analysis of competitiveness conditions to different pathogens and vectors. Moreover, it is still unclear whether non-competitive or beneficial interactions take place among pathogens in insects where multiple infections are observed. Finally, how the plants are implicated in these interactions? Many examples are available on the effects displayed by phytopathogens on the plant processes in favour of insects, such as the promotion of insect attraction to infected hosts, allowing the pathogen spreading (Orlovskis et al., 2015). However, whether pathogens that are capable to modulate their attractiveness could display enhanced competitiveness against horizontally transmitted microbes (including other phytopathogens) is poorly understood. Deep surveys of molecular and cellular machineries of insect-phytopathogen-host plant relations could provide the answers to these issues.

Additional open questions involve the role of bacterial symbionts in plant pathogen competition and spread. Only few examples of interactions between symbionts and pathogens have been described, in spite of the high number of symbiotic bacteria depicted in most of vectors: direct evidences of an interference with the transmission process in the insect or with symptom development in the plant have been provided only for phytoplasmas (Gonella et al., 2018; Iasur-Kruh et al., 2018).

Furthermore, the mechanisms regulating beneficial or hostile exchanges have been only rarely elucidated, and some bacterial pathogens were shown to exhibit mutualistic effects on their vectors, while other caused fitness costs (Hogenhout et al., 2008; Tamborindeguy et al., 2017). An open field for future research is the awareness of whether harmful or beneficial roles are in some way the result of interactions with bacterial symbionts co-inhabiting the same host. A similar evidence of indirect effect on the insect fitness as a consequence of symbiont suppression was observed in virus-transmitting aphids. In the soybean aphid *Aphis glycines* Matsumura, a drop in the concentration of the endosymbiotic *Buchnera* was observed in insects exposed to the beetle-transmitted bean pod mottle virus, resulting in reduced aphid fecundity (Cassone et al., 2015).

Finally, a still unexplored field for vectors of phloem-limited pathogen is the manipulation of symbiotic microbes to drive their interaction with plant pathogens toward antagonistic activities, by means of paratransgenesis. A similar approach was proposed for example for a xylem-restricted pathogenic agent, i.e. the *Xylella fastidiosa* strain causing Pierce disease to grapevine. A bacterium reported as an insect symbiont and an endophyte, *Alcaligenes xyloxydans denitrificans*, was proposed as a candidate agent to be genetically transformed to display anti-*Xylella* molecules (Bextine et al., 2004).

Along with being of certain interest to elucidate biological mechanisms regulating insect-bacteria relationships, the gain of knowledge concerning microbial interactions occurring in insect vectors have important implications for disease epidemiology and control. From the epidemiological point of view, the competition among plant pathogens alters the rates of transmission by vectors, and possibly influences their fitness as well, with a final impact of the spread of diseases on different plants. From the point of view of disease control, the study of microbial interactions in the vectors could provide valuable tools to manage crop infections by altering vector competence via symbiotic control approaches (Alma et al., 2010). Possible strategies include the identification of detrimental effects



1  
2  
3 491 played by symbionts on plant pathogens in the insect, or the selection of new molecular targets to  
4  
5 492 interrupt beneficial interplays among bacteria.  
6  
7

8 493  
9  
10

11 494 **References**  
12  
13

14 495 Alma A, Daffonchio D, Gonella E & Raddadi N (2010) Microbial symbionts of Auchenorrhyncha  
15  
16 496 transmitting phytoplasmas: a resource for symbiotic control of phytoplasmoses. Phytoplasmas -  
17  
18  
19 497 genomes, plant hosts and vectors (ed. by PG Weintraub & P Jones), pp. 272-292. CAB International,  
20  
21 498 Wallingford, UK.  
22

23  
24 499 Alma A, Tedeschi R, Lessio F, Picciau L, Gonella E & Ferracini C (2015) Insect vectors of plant  
25  
26 500 pathogenic Mollicutes in the Euro-Mediterranean region. Phytopathogenic Mollicutes 5: 53-73.  
27

28  
29 501 Arocha-Rosete Y, Kent P, Agrawal V, Hunt D, Hamilton A, Bertaccini A, Scott J, Crosby W &  
30  
31 502 Michelutti R (2011) Preliminary investigations on *Graminella nigrifrons* as a potential vector for  
32  
33 503 phytoplasmas identified at the Canadian Clonal Genebank. Bulletin of Insectology 64: S133-S134.  
34  
35

36  
37 504 Arratia-Castro AA, Santos-Cervantes ME, Arce-Leal AP, Espinoza-Mancillas MG, Rodríguez  
38  
39 505 Negrete EA, Mendez-Lozano J, Arocha-Rosete Y & Leyva-López NE (2016) Detection and  
40  
41 506 quantification of ‘*Candidatus* Phytoplasma asteris’ and ‘*Candidatus* Liberibacter asiaticus’ at early  
42  
43 507 and late stages of Huanglongbing disease development. Canadian Journal of Plant Pathology 38: 411–  
44  
45 508 421.  
46  
47

48  
49 509 Arricau-Bouvery N, Duret S, Dubrana M-P, Batailler B, Desqué D, Béven L, Danet J-L, Monticone  
50  
51 510 M, Bosco D, Malembic-Maher S & Foissac X (2018) Variable membrane protein A of flavescence  
52  
53 511 dorée binds the midgut perimicrovillar membrane of *Euscelidius variegatus* and promotes adhesion  
54  
55 512 to its epithelial cells. Applied and Environmental Microbiology 84: e02487-17.  
56  
57  
58  
59  
60



- Baumann P (2005) Biology of bacteriocyte-associated endosymbionts of plant sap-sucking insects. Annual Review of Microbiology 59: 155–189.
- Bayles BR, Thomas SM, Simmons GS, Grafton-Cardwell EE & Daugherty MP (2017) Spatiotemporal dynamics of the Southern California Asian citrus psyllid (*Diaphorina citri*) invasion. PLoS One 12(3): e0173226.
- Beanland L, Hoy CW, Miller SA & Nault LR (2000) Influence of aster yellows phytoplasma on the fitness of aster leafhopper (Homoptera : Cicadellidae). Annals of the Entomological Society of America 93: 271-276.
- Bendix C & Lewis J (2018) The enemy within: phloem-limited pathogens. Molecular plant pathology 19:238-254.
- Béven L, Duret S, Batailler B, Dubrana M-P, Saillard C, Renaudin J & Arricau-Bouvery N (2012) The repetitive domain of ScARP3d triggers entry of *Spiroplasma citri* into cultured cells of the vector *Circulifer haematoceps*. PLoS ONE 7: e48606.
- Bextine B, Lauzon C, Potter S, Lampe D & Miller TA (2004) Delivery of a genetically marked *Alcaligenes* sp to the glassy-winged sharpshooter for use in a paratransgenic control strategy. Current Microbiology 48: 327-331.
- Bosco D & D'Amelio R (2010) Transmission specificity and competition of multiple phytoplasmas in the same vector. Phytoplasmas - genomes, plant hosts and vectors (ed. by PG Weintraub & P Jones), pp. 293-308. CAB International, Wallingford, UK.
- Bové JM & Garnier M (2002) Phloem-and xylem-restricted plant pathogenic bacteria. Plant Science 163: 1083-1098.
- Bressan A (2014) Emergence and evolution of *Arsenophonus* bacteria as insect-vectored plant pathogens. Infection, Genetics and Evolution 22: 81-90.

- 1  
2  
3 536 Bressan A, Sémétey O, Nusillard B, Clair D & Boudon-Padieu E (2008) Insect vectors (Hemiptera:  
4  
5 537 Cixiidae) and pathogens associated with the disease syndrome “Basses Richesses” of sugar beet in  
6  
7 538 France. *Plant Disease* 92: 113-119.  
9  
10  
11 539 Bressan A, Arneodo J, Simonato M, Haines WP & Boudon-Padieu E (2009a) Characterization and  
12  
13 540 evolution of two bacteriome-inhabiting symbionts in cixiid planthoppers (Hemiptera:  
14  
15 541 Fulgoromorpha: Pentastirini). *Environmental Microbiology* 11: 3265–3279.  
16  
17  
18 542 Bressan A, Sémétey O, Arneodo J, Lherminier J & Boudon-Padieu E (2009b) Vector transmission of  
19  
20 543 a plant pathogenic bacterium in the *Arsenophonus* clade sharing ecological traits with facultative  
21  
22 544 insect endosymbionts. *Phytopathology* 99: 1289–1296.  
23  
24  
25  
26 545 Camerota C, Raddadi N, Pizzinat A, Gonella E, Crotti E, Tedeschi R, Mozes-Daube N, Ember I, Acs  
27  
28 546 Z, Kolber M, Zchori-Fein E, Daffonchio D & Alma A (2012) Incidence of ‘*Candidatus Liberibacter*  
29  
30 547 europaeus’ and phytoplasmas in *Cacopsylla* species (Hemiptera: Psyllidae) and their host/shelter  
31  
32 548 plants. *Phytoparasitica* 40: 213–221.  
33  
34  
35  
36 549 Capoor SP, Rao DG & Viswanath SM (1967) *Diaphorina citri* Kuway, a vector of the greening  
37  
38 550 disease of citrus in India. *Indian Journal of Agricultural Science* 37: 572–576.  
39  
40  
41 551 Caspi-Fluger A & Zchori-Fein E (2010) Do plants and insects share the same symbionts? *Israel*  
42  
43 552 *Journal of Plant Sciences* 58(2): 113-119.  
44  
45  
46 553 Cassone BJ, Redinbaugh MG, Dorrance AE & Michel AP Shifts in *Buchnera aphidicola* density in  
47  
48 554 soybean aphids (*Aphis glycines*) feeding on virus-infected soybean. *Insect Molecular Biology* (2015)  
49  
50 555 24: 422–431.  
51  
52  
53  
54 556 Cicero JM, Fisher TW, Qureshi JA, Stansly PA & Brown JK (2016) Colonization and intrusive  
55  
56 557 invasion of potato psyllid by ‘*Candidatus Liberibacter solanacearum*’. *Phytopathology* 107: 36-49.  
57  
58  
59  
60

- Cooper WR, Garczynski SF, Horton DR, Unruh TR, Beers EH, Shearer W, Richard P & Hilton J (2017) Bacterial endosymbionts of the psyllid *Cacopsylla pyricola* (Hemiptera: Psyllidae) in the Pacific northwestern United States. *Environmental Entomology* 46: 393–402.
- Crotti E, Damiani C, Pajoro M, Gonella E, Rizzi A, Ricci I, Negri I, Scuppa P, Rossi P, Ballarini P, Raddadi N, Marzorati M, Sacchi L, Clementi E, Genchi M, Mandrioli Bandi C, Favia G, Alma A & Daffonchio D (2009) *Asaia*, a versatile acetic acid bacterial symbiont, capable of cross-colonizing insects of phylogenetically distant genera and orders. *Environmental Microbiology* 11: 3252–3264.
- Crotti E, Balloi A, Hamdi C, Sansonno L, Marzorati M, Gonella E, Favia G, Cherif A, Bandi C, Alma A & Daffonchio D (2012) Microbial symbionts: a resource for the management of insect-related problems. *Microbial Biotechnology* 5: 307–317.
- Danet J-L, Foissac X, Zreik L, Salar P, Verdin E, Nourrisseau JG & Garnier M (2003) “*Candidatus* Phlomobacter fragariae” is the prevalent agent of marginal chlorosis of strawberry in French production fields and is transmitted by the planthopper *Cixius wagneri* (China). *Phytopathology* 93: 644–649.
- de Oliveira E, Santos JC, Magalhães PC & Cruz I (2007) Maize bushy stunt phytoplasma transmission by *Dalbulus maidis* is affected by spiroplasma acquisition and environmental conditions. *Bulletin of Insectology* 60: 229–230.
- Duan Y, Zhou L, Hall DG, Li W, Doddapaneni H, Lin H, Liu L, Vahling CM, Gabriel DW, Williams KP, Dickerman A, Sun Y & Gottwald T (2009) Complete genome sequence of citrus Huanglongbing bacterium, '*Candidatus* Liberibacter asiaticus' obtained through metagenomics. *Molecular Plant-Microbe Interactions* 22: 1011–20.
- Duret S, Batailler B, Dubrana M-P, Saillard C, Renaudin J, Béven L & Arricau-Bouvery N (2014) Invasion of insect cells by *Spiroplasma citri* involves spiralin relocalization and lectin/glycoconjugate-type interactions. *Cellular Microbiology* 16: 1119–1132.

- 1  
2  
3 582 Duron O, Bouchon D, Boutin S, Bellamy L, Zhou L & Engelstadter J (2008). The diversity of  
4  
5 583 reproductive parasites among arthropods: *Wolbachia* do not walk alone. BMC Biology 6: 27.  
6  
7  
8 584 Eliautout R, Dubrana M-P, Vincent-Monegat C, Vallier A, Braquart-Varnier C, Poirie M, Saillard C,  
9  
10 585 Heddi A & Arricau-Bouvery N (2016) Immune response and survival of *Circulifer haematoceps* to  
11  
12 586 *Spiroplasma citri* infection requires expression of the gene hexamerin. Developmental and  
13  
14 587 Comparative Immunology 54: 7e19.  
15  
16  
17  
18 588 Fagen JR, Giongo A, Brown CT, Davis-Richardson AG, Gano KA & Triplett EW (2012)  
19  
20 589 Characterization of the relative abundance of the citrus pathogen *Ca. Liberibacter asiaticus* in the  
21  
22 590 microbiome of its insect vector, *Diaphorina citri*, using high throughput 16S rRNA sequencing. Open  
23  
24 591 Microbiology Journal 6, 29–33.  
25  
26  
27  
28 592 Fagen JR, Leonard MT, Coyle JF, McCullough CM, Davis-Richardson AG, Davis MJ & Triplett EW  
29  
30 593 (2014a) *Liberibacter crescens* BT-1T gen. nov.; sp. nov., first cultured member of the *Liberibacter*  
31  
32 594 genus. International Journal of Systematic and Evolutionary Microbiology 64: 2461–2466.  
33  
34  
35 595 Fagen JR, Leonard MT, McCullough CM, Edirisinghe JN, Henry CS, DavisMJ & Triplett EW  
36  
37 596 (2014b) Comparative genomics of cultured and uncultured strains suggests genes essential for free-  
38  
39 597 living growth of *Liberibacter*. PLoS ONE 9: e84469.  
40  
41  
42  
43 598 Freitag JH (1976) Interactions between strains of aster yellows virus in the six-spotted leafhopper  
44  
45 599 *Macrostes fasciifrons*. Phytopathology 57: 1016-1024.  
46  
47  
48 600 Galetto L, Abbà S, Rossi M, Vallino M, Pesando M, Arricau-Bouvery N, Dubrana M-P, Chitarra W,  
49  
50 601 Pegoraro M, Bosco D & Marzachi C (2018) Two phytoplasmas elicit different responses in the insect  
51  
52 602 vector *Euscelidius variegatus* Kirschbaum. Infection and Immunity 86: e00042-18.  
53  
54  
55  
56 603 Galetto L, Marzachi C, Demichelis S & Bosco D (2011) Host plant determines the phytoplasma  
57  
58 604 transmission competence of *Empoasca decipiens* (Hemiptera: Cicadellidae). Journal of Economic  
59  
60 605 Entomology 104: 360-366.

- Galetto L, Nardi M, Saracco P, Bressan A, Marzachi C & Bosco D (2009) Variation in vector competency depends on chrysanthemum yellows phytoplasma distribution within *Euscelidius variegatus* Entomologia Experimentalis et Applicata 131: 200–207.
- Gasparich GE (2010) Spiroplasmas and phytoplasmas: microbes associated with plant hosts. Biologicals 38: 193-203.
- Gatineau F, Jacob N, Vautrin S, Larrue J, Lherminier J, Richard-Molard M & Boudon-Padieu E. (2002) Association with the syndrome "basses richesses" of sugar beet of a phytoplasma and a bacterium-like organism transmitted by a *Pentastiridius* sp. Phytopathology 92: 384-92.
- Gonella E, Negri I, Marzorati M, Mandrioli M, Sacchi L, Pajoro M, Crotti E, Rizzi A, Clementi E, Tedeschi R, Bandi C, Alma A & Daffonchio D (2011) Bacterial endosymbiont localization in *Hyalesthes obsoletus*, the insect vector of "bois noir" in *Vitis vinifera*. Applied and Environmental Microbiology 77: 1423-1435.
- Gonella E, Pajoro M, Marzorati M, Crotti E, Mandrioli M, Pontini M, Bulgari D, Negri I, Sacchi L, Chouaia B, Daffonchio D & Alma A (2015) Plant-mediated interspecific horizontal transmission of an intracellular symbiont in insects. Scientific Reports 5: 15811.
- Gonella E, Crotti E, Mandrioli M, Daffonchio D & Alma A (2018) *Asaia* symbionts interfere with infection by "flavescence dorée" phytoplasma in leafhoppers. Journal of Pest Science 91: 1033–1046.
- Gottwald TR, da Graça JV & Bassanezi RB (2007) Citrus Huanglongbing: the pathogen and its impact. Plant Health Progress DOI:10.1094/PHP-2007-0906-01-RV.
- Haapalainen M (2014) Biology and epidemics of *Candidatus Liberibacter* species, psyllid-transmitted plant pathogenic bacteria. Annals of Applied Biology 165: 172-198.
- Halbert SE & Núñez CA (2004) Distribution of the Asian citrus psyllid, *Diaphorina citri* Kuwayama (Rhynchota: Psyllidae) in the Caribbean basin. Florida Entomologist 87: 401–402.

- 1  
2  
3 629 Hogenhout SA, Oshima K, Ammar E-D, Kakizawa S, Kingdom HN & Namba S (2008)  
4  
5 630 Phytoplasmas: bacteria that manipulate plants and insects. *Molecular Plant Pathology*: 9: 403–423.  
6  
7  
8 631 Iasur-Kruh L, Naor V, Zahavi T, Ballinger MJ, Sharon R, Robinson WE, Perlman SJ & Zchori-Fein  
9  
10 632 E (2017) Bacterial associates of *Hyalesthes obsoletus* (Hemiptera: Cixiidae), the insect vector of bois  
11  
12  
13 633 noir disease, with a focus on cultivable bacteria. *Research in Microbiology* 168: 94–101.  
14  
15  
16 634 Iasur-Kruh L, Zahavi T, Barkai R, Freilich S, Zchori-Fein E & Naor V (2018) *Dyella*-like bacterium  
17  
18 635 isolated from an insect as a potential biocontrol agent against grapevine yellows. *Phytopathology*  
19  
20 636 108: 336-341.  
21  
22  
23 637 Ishii Y, Matsuura Y, Kakizawa S, Nikoh N & Fukatsu T (2013) Diversity of bacterial endosymbionts  
24  
25 638 associated with *Macrostelus* leafhoppers vectoring phytopathogenic phytoplasmas. *Applied and*  
26  
27  
28 639 *Environmental Microbiology* 79: 5013–5022.  
29  
30  
31 640 Jain M, Fleites LA & Gabriel DW (2017) Small *Wolbachia* protein directly represses phage lytic  
32  
33 641 genes in “*Candidatus Liberibacter asiaticus*”. *mSphere* 2: e00171-17.  
34  
35  
36 642 Jing X, Wong AC-N, Chaston JM, Colvin J, McKenzie CL & Douglas AE (2014) The bacterial  
37  
38 643 communities un plant phloem-sap-feeding insects. *Molecular Ecology* 23: 1433-1444.  
39  
40  
41 644 Keremane ML, Ramadugu C, Castaneda A, Diaz JE, Peñaranda EA, Chen J, Duan YP, Halbert SE &  
42  
43  
44 645 Lee RF (2015) Report of *Candidatus Liberibacter caribbeanus*, a new citrus- and psyllid-associated  
45  
46 646 *Liberibacter* from Colombia, South America. American Phytopathological Society Annual Meeting.  
47  
48 647 URL [http://www.apsnet.org/meetings/Documents/2015\\_meeting\\_abstracts/aps2015abO253.htm](http://www.apsnet.org/meetings/Documents/2015_meeting_abstracts/aps2015abO253.htm)  
49  
50  
51 648 Kolora LD, Powell CM, Hunter W, Bextine B & Lauzon CR (2015) Internal extracellular bacteria of  
52  
53 649 *Diaphorina citri* Kuwayama (Hemiptera: Psyllidae), the Asian citrus psyllid. *Current Microbiology*  
54  
55 650 70: 710.  
56  
57  
58 651 Konnerth A, Krczal G & Boonrod K (2016) Immunodominant membrane proteins of phytoplasmas.  
59  
60 652 *Microbiology* 162: 1267–1273.



- 653 Križanac I, Mikec I, Budinščak Ž, Šeruga Musić M & Škori D (2010) Diversity of phytoplasmas  
654 infecting fruit trees and their vectors in Croatia. *Journal of Plant Diseases and Protection*, 117: 206–  
655 213.
- 656 Krstić O, Cvrković T, Mitrović, Radonjić S, Hrnčić S, Toševski I & Jović J (2018) *Wolbachia*  
657 infection in natural populations of *Dictyophara europaea*, an alternative vector of grapevine  
658 Flavescence dorée phytoplasma: effects and interactions. *Annals of Applied Biology* 172: 47-64.
- 659 Kruse A, Fattah-Hosseini S, Saha S, Johnson R, Warwick E, Sturgeon K, Mueller L, MacCoss MJ,  
660 Shatters RG Jr. & Heck MC (2017) Combining 'omics and microscopy to visualize interactions  
661 between the Asian citrus psyllid vector and the Huanglongbing pathogen *Candidatus Liberibacter*  
662 *asiaticus* in the insect gut. *PLoS ONE* 12: e0179531.
- 663 Kruse A, Ramsey JS, Johnson R, Hall DG, MacCoss MJ & Heck M (2018) *Candidatus Liberibacter*  
664 *asiaticus* minimally alters expression of immunity and metabolism proteins in hemolymph of  
665 *Diaphorina citri*, the insect vector Huanglongbing. *Journal of Proteome Research* 17: 2995-3011.
- 666 Kwon MO, Wayadande AC & Fletcher J (1999) *Spiroplasma citri* movement into the intestines and  
667 salivary glands of its leafhopper vector, *Circulifer tenellus*. *Phytopathology* 89: 1144–1151.
- 668 Labroussaa F, Arricau-Bouvery N, Dubrana M-P & Saillard C (2010) Entry of *Spiroplasma citri* into  
669 *Circulifer haematoceps* cells involves interaction between spiroplasma Phosphoglycerate kinase and  
670 leafhopper actin. *Applied and Environmental Microbiology* 76: 1879–1886.
- 671 Labroussaa F, Dubrana M-P, Arricau-Bouvery N, Béven L & Saillard C (2011) Involvement of a  
672 minimal actin-binding region of *Spiroplasma citri* phosphoglycerate kinase in spiroplasma  
673 transmission by its leafhopper vector. *PLoS ONE* 6: e17357.
- 674 Lahav T, Zchori-fein E, Naor V, Freilich S & Iasur-Kruh L (2016) Draft genome sequence of a  
675 *Dyella*-like bacterium from the planthopper *Hyalesthes obsoletus*. *Genome Announcements* 4:  
676 e00686-16.



- 1  
2  
3 677 Landi L, Isidoro N & Riolo P (2013) Natural phytoplasma infection of four phloem-feeding  
4  
5 678 Auchenorrhyncha across vineyard agroecosystems in central–eastern Italy. *Journal of Economic*  
6  
7 679 *Entomology* 106: 604–613.  
8  
9  
10  
11 680 Latorre A & Manzano-Marín A (2017) Dissecting genome reduction and trait loss in insect  
12  
13 681 endosymbionts. *Annals of the New York Academy of Sciences* 1389: 52–57.  
14  
15  
16 682 Lee I-M, Hammond RW, Davis RE & Gundersen DE (1993) Universal amplification and analysis of  
17  
18 683 pathogen 16S rDNA for classification and identification of mycoplasma-like organisms.  
19  
20 684 *Phytopathology* 83: 834–842.  
21  
22  
23  
24 685 Lee I-M, Gundersen-Rindal DE & Bertaccini A (1998a) Phytoplasma: ecology and genomic  
25  
26 686 diversity. *Phytopathology* 88: 1359–1366.  
27  
28  
29 687 Lee I-M, Gundersen-Rindal DE, Davis RE & Bartoszyk I-M (1998b) Revised classification scheme  
30  
31 688 of phytoplasmas based on RFLP analysis of 16S rRNA and ribosomal protein gene sequences.  
32  
33 689 *International Journal of Systematic Bacteriology* 48: 1153–1169.  
34  
35  
36 690 Leonard MT, Fagen JR, Davis-Richardson AG, Davis MJ & Triplett EW (2012) Complete genome  
37  
38 691 sequence of *Liberibacter crescens* BT-1. *Standards in Genomic Sciences* 7: 271–283.  
39  
40  
41  
42 692 Lidor O, Dror O, Hamershlak D, Shoshana N, Belausov E, Zahavi T, Mozes-Daube N, Naor V,  
43  
44 693 Zchori-Fein, E, Iasur-Kruh L & Bahar O (2018) Introduction of a putative biocontrol agent into a  
45  
46 694 range of phytoplasma- and liberibacter-susceptible crop plants. *Pest Management Science* 74: 811–  
47  
48 695 819.  
49  
50  
51  
52 696 Liefting LW, Sutherland PW, Ward LI, Paice KL, Weir BS & Clover GRG (2009) A new ‘*Candidatus*  
53  
54 697 *Liberibacter*’ species associated with diseases of solanaceous crops. *Plant Disease* 93: 208–214.  
55  
56  
57 698 Liu S-L, Liu H-L, Chang S-C & Lin C-P (2011) Phytoplasmas of two 16S rDNA groups are  
58  
59 699 associated with pear decline in Taiwan. *Botanical Studies* 52: 313–320.  
60

- Longone V, González F, Zamorano A, Pino AM, Araya J, Díaz V, Paltrinieri S, Calari A, Bertaccini A, Picciau L, Alma A, & Fiore N (2011) Epidemiological aspects of phytoplasmas in Chilean grapevines *Bulletin of Insectology* 64: S91-S92.
- Mann RS, Pelz-Stelinski K, Hermann SL, Tiwari S & Stelinski LL (2011) Sexual transmission of a plant pathogenic bacterium, *Candidatus Liberibacter asiaticus*, between conspecific insect vectors during mating. *Plos One* 6: e29197.
- Marcone C (2014) Molecular biology and pathogenicity of phytoplasmas. *Annals of Applied Biology* 165: 199-221.
- Marzorati M, Alma A, Sacchi L, Pajoro M, Palermo S, Brusetti L, Raddadi N, Balloi A, Tedeschi R, Clementi E, Corona S, Quaglino F, Bianco PA, Beninati T, Bandi C & Daffonchio D (2006) A novel bacteroides symbiont is localized in *Scaphoideus titanus*, the insect vector of “flavescence dorée” in *Vitis vinifera*. *Applied and Environmental Microbiology* 72: 1467-1475.
- McClean APD & Oberholzer PCJ (1965) Citrus psylla, a vector of the greening disease of sweet orange. *South Africa Journal of Agricultural Science* 8: 297–298.
- McCutcheon JP & Moran NA (2010) Functional convergence in reduced genomes of bacterial symbionts spanning 200 My of Evolution. *Genome Biology and Evolution* 2: 708–718.
- Morris J, Shiller J, Mann R, Smith G, Yen A & Rodoni B (2017). Novel “*Candidatus Liberibacter*” species identified in the Australian eggplant psyllid, *Acizzia solanicola*. *Microbial Biotechnology*, 10(4), 833–844.
- Morrow JL, Hall AAG & Riegler M (2017) Symbionts in waiting: the dynamics of incipient endosymbiont complementation and replacement in minimal bacterial communities of psyllids. *Microbiome* 5: 58.

- 1  
2  
3 722 Nachappa P, Levy J, Pierson E & Tamborindeguy C (2014) Correlation between "*Candidatus*  
4 723 *Liberibacter solanacearum*" infection levels and fecundity in its psyllid vector. *Journal of Invertebrate*  
5  
6 724 *Pathology* 115: 55-61.  
7  
8  
9  
10  
11 725 Nadarasah G & Stavrinides J (2011) Insects as alternative hosts for phytopathogenic bacteria. *FEMS*  
12  
13 726 *Microbiology Reviews* 35: 555-575.  
14  
15  
16 727 Nakabachi A, Ueoka R, Oshima K, Teta R, Mangoni A, Gurgui M, Oldham NJ, van Echten-Deckert  
17  
18 728 G, Okamura K, Yamamoto K, Inoue H, Ohkuma M, Hongoh Y, Miyagishima S, Hattori M, Piel J &  
19  
20 729 Fukatsu T (2013) Defensive bacteriome symbiont with a drastically reduced genome. *Current Biology*  
21  
22 730 23: 1478–1484.  
23  
24  
25  
26 731 Nicolaisen M, Contaldo N, Makarova O, Paltrinieri S & Bertaccini A (2011) Deep amplicon  
27  
28 732 sequencing reveals mixed phytoplasma infection within single grapevine plants. *Bulletin of*  
29  
30 733 *Insectology* 64: S35-S36.  
31  
32  
33 734 Orlovskis Z, Canale MC, Thole V, Pecher P, Lopes JRS & Hogenhout S (2015) Insect-borne plant  
34  
35 735 pathogenic bacteria: getting a ride goes beyond physical contact. *Current Opinion in Insect Science*  
36  
37 736 9: 16-23.  
38  
39  
40  
41 737 Orságová H, Březígová M & Schlesingerová G (2011) Presence of phytoplasmas in hemipterans in  
42  
43 738 Czech vineyards. *Bulletin of Insectology* 64: S119-S120.  
44  
45  
46 739 Palomera V, Bertin S, ROodríguez A, Bosco D, Virla E & Moya-Raygoza G (2012) Is there any  
47  
48 740 genetic variation among native Mexican and Argentinian populations of *Dalbulus maidis* (Hemiptera:  
49  
50 741 *Cicadellidae*)? *Florida Entomologist* (95): 150-155.  
51  
52  
53  
54 742 Perilla-Henao LM & Casteel CL (2016) Vector-borne bacterial plant pathogens: interactions with  
55  
56 743 hemipteran insects and plants. *Frontiers in Plant Science* 7: 1163.  
57  
58  
59  
60

- Perilla-Henao L, Wilson MR & Franco-Lara L (2016) Leafhoppers *Exitianus atratus* and *Amplipcephalus funzaensis* transmit phytoplasmas of groups 16SrI and 16SrVII in Colombia Plant Pathology 65: 1200–1209.
- Prosdocimi EM, Mapelli F, Gonella E, Borin S & Crotti E (2015) Microbial ecology-based methods to characterize the bacterial communities of non-model insects. Journal of Microbiological Methods 119: 110–125.
- Purcell AH (1982) Insect vector relationships with prokaryotic plant pathogens. Annual Review of Phytopathology 20: 397–417.
- Raddadi N, Gonella E, Camerota C, Pizzinat A, Tedeschi R, Crotti E, Mandrioli M, Bianco PA, Daffonchio D & Alma A (2011) ‘*Candidatus Liberibacter europaeus*’ sp. nov. that is associated with and transmitted by the psyllid *Cacopsylla pyri* apparently behaves as an endophyte rather than a pathogen. Environmental Microbiology 13: 414–426.
- Ramsey JS, Johnson RS, Hoki JS, Kruse A, Mahoney J, Hilf ME, Hunter WB, Hall DG, Schroeder FC, MacCoss MJ & Cilia M (2015) Metabolic interplay between the Asian citrus psyllid and its *Profftella* symbiont: an Achilles’ heel of the citrus greening insect vector. PLoS ONE 10: e0140826.
- Ramsey JS, Chavez JD, Johnson R, Hosseinzadeh S, Mahoney JE, Mohr JP, Robison F, Zhong X, Hall DG, MacCoss M, Bruce J & Cilia M (2017) Protein interaction networks at the host–microbe interface in *Diaphorina citri*, the insect vector of the citrus greening pathogen. Royal Society Open Science 4: 160545.
- Rashidi M, D’Amelio R, Galetto L, Marzachi C & Bosco D (2014) Interactive transmission of two phytoplasmas by the vector insect. Annals of Applied Biology 165: 404–413.
- Renaudin J (2006) Sugar metabolism and pathogenicity of *Spiroplasma citri*. Journal of Plant Pathology 88: 129–139.

- 1  
2  
3 767 Richard-Molard M, Garraëssus S, Malatesta G, Valentin P, Fonné G, Gerst M & Grousseau C (1995)  
4  
5 768 Le syndrome des basses richesses – Investigations au champ et tentatives d'identification de l'agent  
6  
7 769 pathogène et du vecteur. Proceedings of the 58th Congrès de l'Institut International de Recherches  
8  
9  
10 770 Betteravières (ed. By F Dijon-Beaune), Brussels, Belgium, pp. 299-309.  
11  
12  
13 771 Rizza S, Pesce A, D'Urso V, Raciti E, Marzachi C & Tessitori M (2016) Transmission of '*Candidatus*  
14  
15 772 *Phytoplasma asteris*' (16SrI) by *Osbornellus horvathi* (Matsumura 1908) co-infected with '*Ca.*  
16  
17 773 *Phytoplasma phoenicium*' (16SrIX). *Phytoparasitica* 44: 491–500.  
18  
19  
20 774 Saldaña MA, Hedge S & Hughes GL (2017) Microbial control of arthropod-borne disease. *Memórias*  
21  
22 do Instituto Oswaldo Cruz 112: 81-93.  
23  
24  
25 776 Saillard C, Vignault JC, Bove JM, Raie A, Tully JG, Williamson DL, Fos A, Garnier M, Gadeau A,  
26  
27 777 Carle P & Whitcomb RF (1987) *Spiroplasma phoeniceum* sp. nov. a new plant-pathogenic species  
28  
29 778 from Syria. *International Journal of Systematic Bacteriology* 37: 106-115.  
30  
31  
32  
33 779 Satta E, Ramirez AS, Paltrinieri S, Contaldo N, Benito P, Poveda JB & Bertaccini A (2016)  
34  
35 780 Simultaneous detection of mixed '*Candidatus* *Phytoplasma asteris*' and '*Ca. Liberibacter*  
36  
37 781 *solanacearum*' infection in carrot. *Phytopathologia Mediterranea* 55: 401–409.  
38  
39  
40  
41 782 Skidmore IH & Hansen AK (2017) The evolutionary development of plant-feeding insects and their  
42  
43 783 nutritional endosymbionts. *Insect Science* 24: 910-928.  
44  
45  
46 784 Swisher KD, Munyaneza JE, Velásquez-Valle R & Mena-Covarrubias J (2018) Detection of pathogens  
47  
48 785 associated with psyllids and leafhoppers in *Capsicum annuum* L. in the Mexican states of Durango,  
49  
50 Zacatecas, and Michoacán. *Plant Disease* 102: 146-153.  
51  
52  
53  
54 787 Tamborindeguy C, Huot OB, Ibanez F & Levy J (2017) The influence of bacteria on multitrophic  
55  
56 788 interactions among plants, psyllids, and pathogen. *Insect Science* 24: 961–974.  
57  
58  
59  
60

- Teixeira DC, Saillard C, Eveillard S, Danet JL, da Costa PI, Ayres AJ & Bové J (2005) '*Candidatus Liberibacter americanus*', associated with citrus huanglongbing (greening disease) in São Paulo State, Brazil. *International Journal of Systematic and Evolutionary Microbiology* 55: 1857-62.
- Terlizzi F, Babini AR, Lanzoni C, Pisi A, Credi R & Foissac X (2007) First report of a gamma 3-proteobacterium associated with diseased strawberries in Italy. *Plant Disease* 91: 1688.
- Thao ML, Moran NA, Abbot P, Brennan EB, Burckhardt DH & Baumann P (2000) Cospeciation of psyllids and their primary prokaryotic endosymbionts. *Applied and Environmental Microbiology* 66: 2898–2905.
- Thinakaran J, Yang XB, Munyaneza JE, Rush CM & Henne DC (2015) Comparative biology and life tables of "*Candidatus Liberibacter Solanacearum*"-infected and -free *Bactericera Cockerelli* (Hemiptera: Triozidae) on potato and silverleaf nightshade. *Annals of the Entomological Society of America* 108: 459-467.
- Thompson S, Fletcher JD, Ziebell H, Beard S, Panda P, Jorgensen N, Fowler SV, Liefting LW, Berry N & Pitman AR (2013) First report of '*Candidatus Liberibacter europaeus*' associated with psyllid infested Scotch broom. *New Disease Reports* 27: 6.
- Wangkeeree J, Miller TA & Hanboonsong Y (2012) Candidates for symbiotic control of sugarcane white leaf disease. *Applied and Environmental Microbiology* 78: 6804–6811.
- Weintraub PG & Beanland L (2006) Insect vectors of phytoplasmas. *Annual Review of Entomology* 51: 91-111.
- Weiss B & Aksoy S (2011) Microbiome influences on insect host vector competence. *Trends in Parasitology* 27: 514-522.

1  
2  
3 810 Wilkes T, Duron O, Darby AC, Hypša V, Nováková E & Hurst GDD (2011). The genus  
4  
5 811 *Arsenophonus*. Manipulative tenants: bacteria associated with arthropods (ed. by E Zchori-Fein & K  
6  
7 812 Bourtzis), pp. 225–244. CRC Press, Boca Raton, FL, USA.  
8  
9  
10  
11 813 Zarei Z, Salehi M, Azami Z, Salari K & Béve L (2017) Stubborn disease in Iran: diversity of  
12  
13 814 *Spiroplasma citri* strains in *Circulifer haematoceps* leafhoppers collected in sesame fields in Fars  
14  
15 815 Province. *Current Microbiology* 74: 239.  
16  
17  
18 816 Zchori-Fein E & Bourtzis K (2011) Manipulative tenants: bacteria associated with arthropods. CRC  
19  
20 817 Press, Boca Raton, FL, USA.  
21  
22  
23 818 Zhao Y & Davis RE (2016) Criteria for phytoplasma 16Sr group/subgroup delineation and the need  
24  
25  
26 819 of a platform for proper registration of new groups and subgroups. *International Journal of Systematic*  
27  
28 820 *and Evolutionary Microbiology*, 66: 2121-2123.  
29  
30  
31 821 Zreik L, Bove J & Garnier M (1998) Phylogenetic characterization of the bacterium-like organism  
32  
33 822 associated with marginal chlorosis of strawberry and proposition of a *Candidatus* taxon for the  
34  
35 823 organism, '*Candidatus* Phlomobacter fragariae'. *International Journal of Systematic Bacteriology* 48:  
36  
37  
38 824 257-261.  
39  
40  
41 825  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60



## Table legends

**Table 1** Multiple bacterial infections in the vectors of phloem-limited pathogens. Only reports showing mixed infections in the same host individual, involving distinct plant disease agents or symbiotic bacteria with phytopathogens, are listed.

**Table 2** Symbiont-pathogen interactions reported in the vectors of phloem-limited plant pathogenic bacteria.

## Figure legends

**Figure 1** Insect symbionts could be useful for controlling the transmission of phloem-limited plant pathogens. Phloem-restricted plant pathogens are indicated as red, purple or violet dots, while microbial symbionts are depicted with other different colours. Microbe movements are indicated with red arrows. Symbiont-mediated control mechanisms of pathogen transmission are listed on the right and corresponding numbers are depicted in gut (in green circles), hemolymph (in orange circles) and salivary glands (inset, in blue circles).

1

2

3 1 TABLES

4

5

6 2

7

8

Vector taxonomic position	Vector family	Vector species	Phytopathogen multiple infection	Symbiont - phytopathogen multiple infection	Reference
Auchenorrhyncha - Fulgoromorpha	Cixiidae	<i>Hyalesthes obsoletus</i> Signoret		' <i>Ca. Sulcia muelleri</i> ' + <i>Wolbachia</i> + ' <i>Ca. Vidania fulgoroidae</i> ' + ' <i>Ca. Purcelliella pentastirinorum</i> ' + 16SrXII phytoplasma	Gonella et al., 2011
		<i>Pentastiridius leporinus</i> L.		' <i>Ca. Sulcia muelleri</i> ' + ' <i>Ca. Purcelliella pentastirinorum</i> ' + <i>Wolbachia</i> + ' <i>Ca. Arsenophonus phytopathogenicus</i> '	Bressan et al., 2009a
		<i>Amplicephalus curtulus</i> Linnavuori & DeLong	Phytoplasmas, groups: 16SrI + 16SrXII		Longone et al., 2011
		<i>Amplicephalus funzaensis</i> Linnavuori	Phytoplasmas, groups: 16SrI + 16SrVII		Perilla-Henao et al., 2016
Auchenorrhyncha - Cicadomorpha	Cicadellidae	<i>Circulifer tenellus</i> (Baker)	16SrVI phytoplasma + <i>S. citri</i>		Lee et al., 1998a Swisher et al., 2018
		<i>Euscelidius variegatus</i> Kirshbaum	Phytoplasmas, groups: 16SrI + 16SrV	bacterium of <i>E. variegatus</i> (BEV) + 16SrI phytoplasma <i>Asaia</i> + 16SrV phytoplasma	Rashidi et al., 2014 Galletto et al., 2009 Gonella et al., 2018
		<i>Euscelis incisus</i> (Kirschbaum)	Phytoplasmas, groups: 16SrI + 16SrIII + ' <i>Ca. Phytoplasma pruni</i> '		Orságová et al., 2011
		<i>Euscelis lineolatus</i> Brulle	Phytoplasmas, groups: 16SrII+16SrXII		Landi et al., 2013
		<i>Exitianus atratus</i> Linnavuori	Phytoplasmas, groups: 16SrI + 16SrVII		Perilla-Henao et al., 2016
		<i>Graminella nigrifrons</i> (Forbes)	Phytoplasmas, groups: 16SrI + 16SrVII 16SrI + 16SrX		Arocha-Rosete et al., 2011
		<i>Macrosteles sexnotatus</i> (Fallén)		' <i>Ca. Sulcia muelleri</i> ' + <i>Nasuia</i> + 16SrI phytoplasma	Ishii et al., 2013
		<i>Macrosteles striifrons</i> Anufriev		' <i>Ca. Sulcia muelleri</i> ' + ' <i>Ca. Nasuia deltocephalinicola</i> ' + 16SrI phytoplasma	Ishii et al., 2013
		<i>Matsumuratettix hiroglyphicus</i> (Matsumura)		Bacterium associated with <i>M. hiroglyphicus</i> (BAMH) ( <i>Nasuia</i> ) + ' <i>Ca. Sulcia muelleri</i> ' + 16SrXI phytoplasma	Wangkeeree et al., 2012
		<i>Osbornellus horvathi</i> Matsumura	' <i>Ca. Phytoplasma asteris</i> ' + ' <i>Ca. Phytoplasma phoenicium</i> '		Rizza et al., 2016

		<i>Paratanus exitiosus</i> (Beamer)	Phytoplasmas, groups: 16SrII + 16SrVII + 16SrXII	Longone et al., 2011	
		<i>Recilia dorsalis</i> Motschulsky	BAMH + ‘ <i>Ca. Sulcia muelleri</i> ’ +16SrXI phytoplasma	Wangkeeree et al., 2012	
		<i>Recilia</i> sp. nr. <i>vetus</i>	BAMH + ‘ <i>Ca. Sulcia muelleri</i> ’ +16SrXI phytoplasma	Wangkeeree et al., 2012	
		<i>Scaphoideus titanus</i> Ball	‘ <i>Ca. Cardinium hertigii</i> ’ + 16SrV phytoplasma	Marzorati et al., 2006	
		<i>Cacopsylla chinensis</i> (Yang & Li)	Phytoplasmas, groups: 16SrII + 16SrX	Liu et al., 2011	
		<i>Cacopsylla melanoneura</i> (Förster)	CLeu + ‘ <i>Ca. Phytoplasma mali</i> ’	Camerota et al., 2012	
	Psyllidae	<i>Cacopsylla pyri</i> L.	Phytoplasmas, groups: 16SrI + 16SrXII 16SrX + 16SrXII 16SrI + 16SrX	CLeu + ‘ <i>Ca. Carsonella ruddii</i> ’ + <i>Arsenophonus</i> + <i>Ralstonia</i> + ‘ <i>Ca. Phytoplasma pyri</i> ’	Križanac et al., 2010 Raddadi et al., 2011 Camerota et al., 2012
		<i>Cacopsylla pyricola</i> Förster		<i>Arsenophonus</i> + ‘ <i>Ca. Phytoplasma pyri</i> ’ ‘ <i>Ca. Carsonella ruddii</i> ’ + ‘ <i>Ca. Proffittella aramtura</i> ’ + <i>Wolbachia</i> + CLas	Cooper et al., 2017 Kruse et al., 2017 Ramsey et al., 2017
		<i>Diaphorina citri</i> Kuwayama		<i>Ralstonia</i> + CLas	Kolora et al., 2015
		<i>Bactericera cockerelli</i> (Sulc)		<i>Erwinia</i> sp. + <i>Wolbachia</i> + <i>Staphylococcus</i> sp. + <i>Enterococcus</i> sp. + CLso	Kolora et al., 2015
	Triozidae	<i>Bactericera trigonica</i> Hodkinson	CLso + phytoplasmas, (group 16SrVI+ 16SrI)		Swisher et al., 2018

**Table 2** Symbiont-pathogen interactions reported in the vectors of phloem-limited plant pathogenic bacteria.

Insect	Phytopathogen	Symbiont	Interaction	Reference
<i>Diaphorina citri</i>	CLas	' <i>Ca. Proffittella armatura</i> '	Upregulation of genes involved in biosynthesis of diaphorin polyketide.	Ramsey et al., 2015
<i>Hyaletbes obsoletus</i>	16SrXII phytoplasma	<i>Wolbachia</i> <i>Dyella</i> -like bacterium (DLB)	Positive correlation Reduction of phytoplasma-related symptoms in grapevine	Fagen et al., 2012 lasur-Kruh et al., 2018
<i>Euscelidius variegatus</i>	16SrV phytoplasma	<i>Asaia</i> sp.	Reduced phytoplasma acquisition in <i>Asaia</i> -infected individuals	Gonella et al., 2018
<i>Matsumuratettix hiroglyphicus</i> <i>Recilia dorsalis</i> <i>Recilia</i> sp. nr. <i>vetus</i>	phytoplasmas	Bacterium associated with <i>M. hiroglyphicus</i> (BAMH) ( <i>Nasuia</i> )	BAHM suggested to be required for successful phytoplasma transmission	Wangkeeree et al., 2012
<i>Dyctiophara europaea</i>	16SrV phytoplasma	<i>Wolbachia</i>	Mutual exclusion	Krstić et al., 2018